IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant:

Osamu OKUDA et al.

Title:

METHODS FOR TREATING INTERLEUKIN-6

RELATED DISEASES

Appl. No.:

Unassigned

Filing Date: Herewith

Examiner:

Unassigned

Art Unit:

Unassigned

INFORMATION DISCLOSURE STATEMENT UNDER 37 CFR §1.56

Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir:

Submitted herewith on Form PTO/SB/08 is a listing of documents known to Applicants in order to comply with Applicants' duty of disclosure pursuant to 37 CFR §1.56.

The submission of any document herewith, which is not a statutory bar, is not intended as an admission that such document constitutes prior art against the claims of the present application or that such document is considered material to patentability as defined in 37 CFR §1.56(b). Applicants do not waive any rights to take any action which would be appropriate to antedate or otherwise remove as a competent reference any document which is determined to be a prima facie art reference against the claims of the present application.

TIMING OF THE DISCLOSURE

The listed documents are being submitted in compliance with 37 CFR §1.97(b), within three (3) months of the date of entry of the national stage as set forth in 37 CFR §1.491.

RELEVANCE OF EACH DOCUMENT

All of the documents listed on the attached PTO/SB/08 were cited as being relevant during the prosecution of the corresponding International application. Copies of the documents are not being provided since copies should have been provided directly by WIPO under an exchange program between the PTO, the EPO and the JPO. A copy of the International Search Report is attached setting forth the portion of each document considered relevant by the examiner.

Applicants respectfully request that each listed document be considered by the Examiner and be made of record in the present application and that an initialed copy of Form PTO/SB/08 be returned in accordance with MPEP §609.

The Commissioner is hereby authorized to charge any additional fees which may be required regarding this application under 37 CFR §§ 1.16-1.17, or credit any overpayment, to Deposit Account No. 19-0741. Should no proper payment be enclosed herewith, as by a check being in the wrong amount, unsigned, post-dated, otherwise improper or informal or even entirely missing, the Commissioner is authorized to charge the unpaid amount to Deposit Account No. 19-0741.

Respectfully submitted,

Date October 24, 2005

FOLEY & LARDNER LLP Customer Number: 22428

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Sheet	•	1	of	2	Attorney Docket Number	053466-0409			

			U.S. PATENT DOCUMENTS	3		
 0:1-	U.S. Patent Document		Name of Botonton on Applicant of	Date of Publication of	Pages, Columns, Lines, Where Relevant	
 Cite No. ¹	Number	Kind Code ² (if known)	Name of Patentee or Applicant of Cited Document	Cited Document MM-DD-YYYY	Passages or Relevant Figures Appear	
 A1	5,210,075		Scholz et al.	05/11/1993		
A2	6,270,766	B1	Feldman et al.	08/07/2001		
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Examiner Initials*	Cite No. ¹	Fo Office ³	oreign Patent Document Number ⁴ Kind Co	ode ⁵	Name of Patentee or Applicant of Cited Documents	Date of Publication of Cited Document MM-DD-YYYY	Pages, Columns, Lines, Where Relevant Passages or Relevant Figures Appear	T ⁶
	A3	EP	1 074 268	A1	Chugai Seiyaku Kabushiki Kaisha and Tadamitsu KISHIMOTO	02/07/2001		
	A4	PCT	WO 97/10338		Chiron Corporation	03/20/1997		
	A5	PCT	WO 99/64070		Ophidian Pharmaceuticals, Inc.	12/16/1999		
4,	A6	PCT	WO 2004/039826	A1	Centocor, Inc.	05/13/2004		

		NON PATENT LITERATURE DOCUMENTS			
Examiner Initials*	Cite No. Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.) date, page(s), volume-issue numbe publisher, city and/or country where published.				
	A7	BAERT, Filip et al, "Influence of Immunogenicity on the Long-Term Efficacy of Infliximab in Crohn's Disease", NEW ENGLAND JOURNAL OF MEDICINE, February 2003, Vol. 348, No. 7, pgs. 601-608.			
	A8	CHOY, E. H. S. et al, "Therapeutic Benefit of Blocking Interleukin-6 Activity With an Anti-Interleukin-6 Receptor Monoclonal Antibody in Rheumatoid Arthritis: A Randomized, Double-Blind, Placebo-Controlled, Dose-Escalation Trial", ARTHRITIS AND RHEUMATISM, December 2002, Vol. 46, No. 12, pgs. 3143-3150.			
	A9	ITO, Hiroaki et al., "A Pilot Randomized Trial of a Human Anti-Interleukin-6 Receptor Monoclonal Antibody in Active Crohn's Disease", GASTROENTEROLOGY, April 2004, Vol. 126,. No. 4, pgs. 989-996.			
	A10	MAINI & CHARISMA STUDY GROUP, "A Double-Blind, Parallel Group, Controlled, Dose Ranging Study of the Safety, Tolerability, Phamacokinetics and Efficacy of Repeat Doses of MRA Given Alone or in Combination With Methotrexate in Patients With Rheumatoid Arthritis", ABSTRACT OF PRESENTATION AT EULAR, June 2003, 2 pages.			

Examiner Signature		Date Conside	ered	

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	Date Submitte	ed: October 24, 2005	Group Art Unit	Unassigned				
	(use as many	sheets as necessary)	Examiner Name	Unassigned				
Sheet	2	of 2	Attorney Docket Number	053466-0409				

		NON PATENT LITERATURE DOCUMENTS	
Examiner Initials*	Cite No.1	Include name of the author (in CAPITAL LETTERS), title of the article (when appropriate), title of the item (book, magazine, journal, serial, symposium, catalog, etc.) date, page(s), volume-issue number(s), publisher, city and/or country where published.	T⁵
	A11	MAINI, Ravinder et al., "Therapeutic Efficacy of Multple Intravenous Infusions of Anti-Tumor Necrosis Factor α Monoclonal Antibody Combined with Low-Dose Weekly Methotrexate in Rheumatoid Arthritis", ARTHRITIS AND RHEUMATISM, September 1998, Vol. 41, No. 9, pgs. 1552-1563.	
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	A14	NISHIMOTO, Norihito et al, "The Long-term Safety and Efficacy of Humanized Anti-Interleukin-6 Receptor Monoclonal Antibody, MRA in Multicentric Castelman's Disease", DATABASE BIOSIS "Online BIOSCIENCES INFORMATION SERVICE, November 2003, 1 page.	
	A15	WAGNER, C. L., et al., "Consequences of Immunogenicity to the Therepeutic Monoclonal Antibodies ReoPro® and Remicade®", IMMUNOGENICITY OF THERAPEUTIC BIOLOGICAL PRODUCTS, 2003, pgs. 37-53.	

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